		•	1 = 1	4019 Rec'd PCT/PTO 0 1 JUN 2001
FORM I	TO-1390	(Modified) U.S. DEPARTMENT	OF COMMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY'S DOCKET NUMBER
(REV 11	-98) TR	ANSMITTAL LETTER	TO THE UNITED STATES	JMYT-245US
ĺ				U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR 1.5)
l		DESIGNATED/ELECTE		To Be Assigned 9 / 857116
ᆫ		CONCERNING A FILIN	INTERNATIONAL FILING DATE	PRIORITY DATE CLAIMED
INTE		IONAL APPLICATION NO. PCT/GB99/03958	29 November 1999 (29.11.99)	01 December 1998 (01.12.98)
TITLE	OFR	NVENTION		
REA	CT0	R AND PROCESS FOR RE	MOVAL OF CARBON MONOXIDE	FROM HYDROGEN
L				
		r(s) FOR DO/EO/US		
		eter Geoffrey , John Frederick		
			ates Designated/Elected Office (DO/EO/US)	the following items and other information:
Appl	cant l			
1.	\boxtimes		tems concerning a filing under 35 U.S.C. 37	
2.		This is a SECOND or SUBSEQ	QUENT submission of items concerning a fil	ing under 35 U.S.C. 371.
3.	\boxtimes	examination until the expiration	gin national examination procedures (35 U.S. of the applicable time limit set in 35 U.S.C	. 3/1(b) and FC1 Articles 22 and 3/(1).
4.	\boxtimes	A proper Demand for Internation	nal Preliminary Examination was made by the	ne 19th month from the earliest claimed priority date.
5.	\boxtimes		lication as filed (35 U.S.C. 371 (c) (2))	
L.		a. 🛛 is transmitted herewith	(required only if not transmitted by the Int	ernational Bureau).
- 100 m		b. has been transmitted b	y the International Bureau.	
100		c. is not required, as the	application was filed in the United States Re	eceiving Office (RO/US).
6.		A translation of the Internationa	l Application into English (35 U.S.C. 371(c)(2)).
÷7.	\boxtimes	A copy of the International Sear		
	\boxtimes	Amendments to the claims of th	e International Application under PCT Artic	tle 19 (35 U.S.C. 371 (c)(3))
juk.		a. are transmitted herewi	th (required only if not transmitted by the Ir	sternational Bureau).
121			by the International Bureau.	
8		c. have not been made; h	owever, the time limit for making such ame	ndments has NOT expired.
500		d. have not been made ar	nd will not be made.	
140		A translation of the amendment	s to the claims under PCT Article 19 (35 U.	S.C. 371(c)(3)).
UPPE	\bowtie	An oath or declaration of the in-	ventor(s) (35 U.S.C. 371 (c)(4)). (UNEXE	CUTED)
41.	×	A copy of the International Prel	iminary Examination Report (PCT/IPEA/40	9).
12.		A translation of the annexes to (35 U.S.C. 371 (c)(5)).	the International Preliminary Examination R	teport under PCT Article 36
1	tems	13 to 20 below concern docume	nt(s) or information included:	
13.		An Information Disclosure Stat	tement under 37 CFR 1.97 and 1.98.	
14.			cording. A separate cover sheet in complian	ace with 37 CFR 3.28 and 3.31 is included.
15.	×	A FIRST preliminary amendm		
16.		A SECOND or SUBSEQUEN		
17.		A substitute specification.	•	
18.		A change of power of attorney	and/or address letter.	

19. ⊠ Certificate of Mailing by Express Mail
 20. □ Other items or information:

Page 1 of 2

U.S. APPLICATION	(IDKROWN ED 87 CFR 1.5)	INTERNATIONAL A	PPLICATI	ON NO.		ATTORNEY'S	DOCKET NUMBER
To Be	ASighed I I D	PCT/GE	399/0395	58		JMY	T-245US
21. The follow	ving fees are submitted:.					CALCULATIONS	PTO USE ONLY
	FEE (37 CFR 1.492 (a) (1) -						
international se	ational preliminary examinatio earch fee (37 CFR 1.445(a)(2) aal Search Report not prepared	paid to USPTO	nor	\$97	0.00		
☑ International property of the Section International Property of	reliminary examination fee (37 ternation Search Report prepar	7 CFR 1.482) not paid red by the EPO or JPO	to	\$84	10.00		
but internation	reliminary examination fee (3° al search fee (37 CFR 1.445(a	(2) paid to USPTO.		O \$69	0.00		
but all claims	reliminary examination fee pa did not satisfy provisions of PC	CT Article 33(1)-(4)		\$67	70.00		
☐ International p and all claims	reliminary examination fee pa satisfied provisions of PCT Ar	ticle 33(1)-(4)			6.00		
	ENTER APPROPRI					\$860.00	
Surcharge of \$130.00 months from the earlie	for furnishing the oath or declar est claimed priority date (37 C	CFR 1.492 (e)).	□ 20			\$0.00	
CLAIMS	NUMBER FILED	NUMBER EXT	RA	RATI			
Total claims	14 - 20 =	0		x \$18.0		\$0.00	
Independent claims	2 - 3=	0		x \$80.0)0	\$0.00	
Multiple Dependent 0	Claims (check if applicable).					\$0.00	
		ABOVE CALC			=	\$860.00	
Reduction of 1/2 for fi must also be filed (No	iling by small entity, if applicate 37 CFR 1.9, 1.27, 1.28) (c	able. Verified Small E heck if applicable).	intity Sta	tement		\$0.00	
T)			SUB'	TOTAL	=	\$860.00	
Processing fee of \$130 months from the earlie	0.00 for furnishing the English est claimed priority date (37 C	translation later than CFR 1.492 (f)).	□ 2	0 🗆 3	+	\$0.00	
<u> </u>		TOTAL NAT	IONAI	LFEE	=	\$860.00	
Fee for recording the	enclosed assignment (37 CFR propriate cover sheet (37 CFR					\$0.00	
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انما: Please charge	e amount of \$860.00 e my Deposit Account No. copy of this sheet is enclosed.	to cover the above f	ees is en			to cover the abo	ve fees.
☑ The Commiss	sioner is hereby authorized to	charge any fees which	may be re	equired, or	credit a	any overpayment	
to Deposit Ac	ecount No. 18-0350	A duplicate copy of the	is sheet i	s enclosed.			
NOTE: Where an a 1.137(a) or (b)) must	ppropriate time limit under t be filed and granted to rest	37 CFR 1.494 or 1.49 ore the application to	5 has not	t been met, status.	a peti	ition to revive (37 C	FR
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Christopher R. Lew	ris			SIGNAT	TIRE	<u> </u>	
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Suite 301				Christo	pher	R. Lewis	
One Westlakes, Ber P.O. Box 980	- wyn			NAME			
Valley Forge, PA 1	9482-0980	:		36,201			
m.l. b (620) /	07.0700			REGIST	RATIC	N NUMBER	
Telephone: (610) 4 Facsimile: (610) 4				June 1,	2001		
(020)							
				DATE			

09/857116

CERTIFICATE OF I	MAIL" (37 CFR 1.10)	JMYT-245US	
Serial No. To Be Assigned	Examiner	Group Art Unit	
vention: REACTOR	AND PROCESS FOR REMOVAL	L OF CARBON MONOXIDE F	ROM HYDROGEN
I hereby certify that th	e following correspondence:		
U.S. National Phase Ap	oplication, w/Form PTO-1390 and	all of the documents listed there	ein,
	(Identify type	of correspondence)	
June 1, (Date		Kathleen Li (Typed or Printed Name of Person M	
μ. Τ		(Signature of Person Mailing	Correspondence)
		EL73229182	
		("Express Mail" Mailing L	abet Number)
	Note: Each paper must ha	ve its own certificate of mailing.	

JC18 Rec'd PCT/PTO 0 1 JUN 2001 PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:

Peter Geoffrey Gray et al.

: Art Unit:

Application No.:

To Be Assigned

: Examiner:

Filed:

Herewith

FOR: REACTOR AND PROCESS FOR

:

REMOVAL OF CARBON MONOXIDE FROM HYDROGEN

.

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington, DC 20231

SIR:

Prior to examination, please amend the above-identified application as follows.

IN THE SPECIFICATION:

Please replace the paragraph beginning at page 3, line 9, with the following:

An essentially cylindrical metal vessel, made by pressing two halves and seam-welding, has a main gas inlet, 2, for reformate gas, and an outlet, 3, for cleaned gas. The vessel has a number of annular reaction zones, 4, each consisting of an annular wound corrugated catalyst support. The vessel further has an internal water jacket, 5, between a spacer and the shell and this is supplied with cooling water by means of a water circuit comprising a circulating pump, 6, and a heat exchanger, 7. The heat

exchanger is desirably used to recover heat in an integrated system e.g. in an integrated combined heat and power system. The vessel further is provided with four air inlets, 8, each providing pre-determined (for a steady-state operation) levels of air (that is reaction oxygen; air is generally suitable). Although not preferred, controlled amounts of oxygen, according to one or more feedback or feed forward controls detecting gas flow levels and CO levels may be used. Downstream of each air inlet is a gas mixing zone, 9a, 9b, 9c and 9d. Zone 9a is suitably a static mixer, but zones 9b, 9c and 9d are preferably alternating annular mixing vanes, 10, and discs, 11, having gas flow holes, as illustrated in Figs 2 and 3. It should be noted that for simplicity only four vanes and four holes are shown, but more, e.g. six or eight, may be used. Such vanes and discs may be manufactured by pressing to form a sleeve which may be fitted and brazed or welded onto the jacket 5 with the intention that each gas mixing zone also serves to remove heat from the reaction gases leaving each reaction zone. Although the schematic drawing of the reactor shows each reaction zone and gas mixing zone of the same size, these may be designed to differ according to the amount of heat to be removed. Additionally, each reaction zone may incorporate different and/or different loadings, of catalyst. Each metal catalyst support monolith suitably has 300-400 cells per sq.in. and is coated with a washcoat comprising ~36wt% alumina solids of D50 of approx 5µm and D90 of approx 16µm, to deposit approx 3g/m³. After drying and firing, the coated monolith was impregnated with a platinum salt solution to result in 5% by wt of Pt deposited on the alumina. The resulting coated monolith was again dried and fired before being assembled into the reactor.

Please replace the paragraph beginning at page 5, line 1, with the following:

The detailed operating conditions and other experimental detail for one typical run of the above reactor now follow. It was remarkable that the pressure drop across the whole of the test reactor was about the

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measurement limit of the gauges being used, and was of the order of 1.5-3mbar (150-300Pa), at the two throughputs tested. This indicates that the reactor is very efficient, compact and requires very little energy to operate.

IN THE CLAIMS:

Please replace claims 1, 3, 4, 6-8, 10, 11, and 14 with the following amended claims:

- A catalytic selective oxidation reactor, comprising a
 cylindrical reactor vessel with counter current cooling means and at least one
 stage, each stage being provided with an inlet for a first feedstock, and inlet
 for a second feedstock, gas mixing means and a catalytic reaction zone,
 wherein the reaction zone comprises a selective oxidation catalyst deposited
 upon a metal support.
- 1 3. A reactor according to claim 1, wherein the metal 2 support is a metal monolith.
 - 4. A reactor according to claim 1, wherein the cooling means comprises a central jacket mounted within the reactor.
- 6. A reactor according to claim 4, wherein the central jacket is connected via a coolant circuit to a circulating pump.
- 1 7. A reactor according to claim 1, wherein the catalytic 2 reaction zone(s) is/are annular and surround a central cooling means.
- 1 8. A reactor according to claim 1, having from 2 to 10 2 stages.
- 1 10. A reactor according to claim 1, wherein the gas mixing means comprise annular mixing vanes or discs.

- 1 11. A reactor according to claim 1, connected to the output
- 2 from a reformer such that CO is removed from the output gas to a level
- 3 where the output from the reactor may be fed to a fuel cell.
- 1 14. A process according to claim 12, providing hydrogen to 2 a fuel cell.

Respectfully submitted,

Christopher R. Lewis, Reg. No. 36,201 Attorney for Applicants

CRI /lrb

Dated: June 1, 2001

Suite 301 One Westlakes, Berwyn P.O. Box 980 Valley Forge, PA 19482-0980 (610) 407-0700

The Assistant Commissioner for Patents is hereby authorized to charge payment to Deposit Account No. 18-0350 of any fees associated with this communication.

EXPRESS MAIL Mailing Label No.: EL732291825US
Date of Deposit: June 1, 2001

I hereby certify that this paper and fee are being deposited, under 37 C.F.R. § 1.10 and with sufficient postage, using the "Express Mail Post Office to Addressee" service of the United States Postal Service on the date indicated above and that the deposit is addressed to the Assistant Commissioner for Patents. Washindton, D.C. 20231, D.C. 20231,

Kathleen Libby

VERSION WITH MARKINGS TO SHOW CHANGES MADE

IN THE SPECIFICATION:

Specification at page 3, line 9:

An essentially cylindrical metal vessel, made by pressing two halves and seam-welding, has a main gas inlet, 2, for reformate gas, and an outlet, 3, for cleaned gas. The vessel has a number of annular reaction zones, 4, each consisting of an annular wound corrugated catalyst support. The vessel further has an internal water jacket, 5, between a spacer and the shell and this is supplied with cooling water by means of a water circuit comprising a circulating pump, 6, and a heat exchanger, 7. The heat exchanger is desirably used to recover heat in an integrated system e.g. in an integrated combined heat and power system. The vessel further is provided with four air inlets, 8, each providing pre-determined (for a steady-state operation) levels of air (that is reaction oxygen; air is generally suitable). Although not preferred, controlled amounts of oxygen, according to one or more feedback or feed forward controls detecting gas flow levels and CO levels may be used. Downstream of each air inlet is a gas mixing zone, 9a, 9b, 9c and 9d. Zone 9a is suitably a static mixer, but zones 9b, 9c and 9d are preferably alternating annular mixing vanes, 10, and discs, 11, having gas flow holes, as illustrated in Figs 2 and 3. It should be noted that for simplicity only four vanes and four holes are shown, but more, e.g. six or eight, may be used. Such vanes and discs may be manufactured by pressing to form a sleeve which may be fitted and brazed or welded onto the jacket 5 with the intention that each gas mixing zone also serves to remove-hear heat from the reaction gases leaving each reaction zone. Although the schematic drawing of the reactor shows each reaction zone and gas mixing zone of the same size, these may be designed to differ according to the amount of heat to be removed. Additionally, each reaction zone may incorporate different

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and/or different loadings, of catalyst. Each metal catalyst support monolith suitably has 300-400 cells per sq.in. and is coated with a washcoat comprising ~36wt% alumina solids of D50 of approx 5µm and D90 of approx 16µm, to deposit approx 3g/m³. After drying and firing, the coated monolith was impregnated with a platinum salt solution to result in 5% by wt of Pt deposited on the alumina. The resulting coated monolith was again dried and fired before being assembled into the reactor.

Specification at page 5, line 1:

The detailed operating conditions and other experimental detail for one typical run of the above reactor now follow. It was remarkable that the pressure drop across the whole of the est test reactor was about the measurement limit of the gauges being used, and was of the order of 1.5-3mbar (150-300Pa), at the two throughputs tested. This indicates that the reactor is very efficient, compact and requires very little energy to operate.

IN THE CLAIMS:

- 1. A catalytic selective oxidation reactor, comprising a cylindrical reactor vessel-(1) with counter current cooling means-(5,6,7) and at least one stage, each stage being provided with an inlet for a first feedstock (2), and inlet for a second feedstock-(8), gas mixing means-(9) and a catalytic reaction zone-(4), wherein the reaction zone comprises a selective oxidation catalyst deposited upon a metal support.
- 1 3. A reactor according to any one of the preceding claims 2 claim 1, wherein the metal support is a metal monolith.
- 4. A reactor according to any one of the preceding claims

 claim 1, wherein the cooling means comprises a central jacket mounted

 within the reactor.

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- A reactor according to claim 4-or 5, wherein the central 6. jacket is connected via a coolant circuit to a circulating pump. 2
- A reactor according to any one of the preceding claims 1 claim 1, wherein the catalytic reaction zone(s) is/are annular and surround a 2 central cooling means. 3
- A reactor according to any one of the preceding claims claim 1, having from 2 to 10 stages. 2
- A reactor according to any one of the preceding claims 10. 1 claim 1, wherein the gas mixing means comprise annular mixing vanes or 2 3 discs.
- A reactor according to any one of the preceding claims 11. 1 claim 1, connected to the output from a reformer such that CO is removed 2 from the output gas to a level where the output from the reactor may be fed 3 to a fuel cell. 4
 - A process according to claim 12-or-13, providing hydrogen to a fuel cell.

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REACTOR AND PROCESS FOR REMOVAL OF CARBON MONOXIDE FROM HYDROGEN

The present invention concerns an improved reactor, more particularly it concerns a reactor particularly adapted for and suitable for certain selective oxidation reactions.

We have previously developed certain selective oxidation processes for the removal of residual carbon monoxide in hydrogen-containing gases intended as feedstock for fuel cells. Such processes are disclosed and claimed in for example, PCT/GB98/02873. It has become well established that hydrogen-fuelled fuel cells require very low proportions of carbon monoxide, which acts as a catalyst poison and degrades performance of the fuel cell. Desirably, the amount of CO is less than 10 ppm. If a liquid fuel is reformed to form a hydrogen-containing fuel, levels of CO are produced which may vary about 0.5-5mol%, so that it becomes necessary to attenuate such CO levels. As mentioned in the above International Application, catalytic selective oxidation reactions are highly exothermic with the result that increasing reaction temperatures lead to increasing loss of valuable hydrogen due to competing reactions.

There remains a need for a catalytic selective oxidation reactor which permits good temperature control yet is simple, effective and inexpensive to manufacture. Although hydrogen "cleaned-up" by the reactor of the present invention may be used as fuel cell fuel, there are many other uses for such hydrogen. Also, although the reactor is particularly suitable for the cleaning up of reformate (which may be from methanol, other oxygenated fuels, gasoline or other hydrocarbon fuels), it is not limited to such use. Accordingly, the present invention provides a reactor for catalytic selective oxidation, comprising a reactor vessel with counter current cooling and at least one stage, each stage provided with an inlet for a first feedstock, e.g. CO contaminated hydrogen, an inlet for a second feedstock, e.g. oxygen-containing gas, gas mixing means and a catalytic reaction zone, preferably comprising a selective oxidation catalyst deposited on a substrate having good heat conduction properties.

The invention also provides a process for the removal of quantities of CO from a

35 hydrogen feedstock by partial oxidation, comprising at least one step of admixing said

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feedstock with a quantity of oxygen, passing the resulting mixture over a selective oxidation catalyst in a selective oxidation zone and recovering a hydrogen product with reduced content of CO, whilst simultaneously cooling the selective oxidation zone, preferably to maintain the selective oxidation zone at a temperature in the range 100 to 250°C, by countercurrent coolant flow.

The reactor, in its most preferred embodiment, has four stages and this is effective to reduce a reformate having a CO content of 1 to 2% to below 10ppm. In other embodiments, there may be 2 to 10 stages.

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The reactor is suitably in the form of a vessel in which the reaction zones are generally annular in shape, and provided with internal and/or external liquid cooling. Cooling is preferably provided by internal cooling using water, which is suitably under pressure to prevent boiling at the temperatures generated. Other liquid coolants, or forced air cooling may be used if desired. According to specific reactor designs, heat removal may be enhanced by increasing heat exchange surface areas by corrugation or other established means. Heat removal may be altered by altering the flow of coolant and/or altering the quantity of heat removed from the coolant by means of a heat exchanger.

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Each reaction zone suitably comprises a metallic catalyst support ("monolith") of generally honeycomb construction, as is in common use in automobile exhaust gas catalysis. These may have 50 to 1200 cells per square in cell density, preferably 200 to 600 cpsi, and may be of a variety of types of stainless steel. In general, the present invention does not result in the reaction stages reaching the higher temperatures generally met with in exhaust gas catalysis (e.g. 500-900°C), but the material should be capable of resisting temperature excursions. An alternative reaction zone design utilises what has become known as "static mixers" which combine a very high degree of gas agitation and mixing, resulting in non-laminar flow, with relatively low pressure drop. Such static mixers may be manufactured from a variety of metals and are commercially available. However, it is presently preferred to use a honeycomb monolith.

The reaction zone desirably carries a selective oxidation catalyst comprising a platinum group metal, especially comprising platinum and/or rhodium, carried on a high

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surface area metal oxide support, such as alumina. The metallic catalyst support may be coated by generally known methods with a slurry of oxide support, dried and fired, and then impregnated with the catalytically active components, before a final firing. Such procedures are in themselves known, and do not form part of this invention.

The invention will now be described with reference to the accompanying drawing which shows a cross-section of one embodiment of the present invention.

An essentially cylindrical metal vessel, made by pressing two halves and seam-welding, has a main gas inlet, 2, for reformate gas, and an outlet, 3, for cleaned gas. The vessel has a number of annular reaction zones, 4, each consisting of an annular wound corrugated catalyst support. The vessel further has an internal water jacket, 5, between a spacer and the shell and this is supplied with cooling water by means of a water circuit comprising a circulating pump, 6, and a heat exchanger, 7. The heat exchanger is desirably used to recover heat in an integrated system e.g. in an integrated combined heat and power system. The vessel further is provided with four air inlets, 8, each providing pre-determined (for a steady-state operation) levels of air (that is reaction oxygen; air is generally suitable). Although not preferred, controlled amounts of oxygen, according to one or more feedback or feed forward controls detecting gas flow levels and CO levels may be used. Downstream of each air inlet is a gas mixing zone, 9a, 9b, 9c and 9d. Zone 9a is suitably a static mixer, but zones 9b, 9c and 9d are preferably alternating annular mixing vanes, 10, and discs, 11, having gas flow holes, as illustrated in Figs 2 and 3. It should be noted that for simplicity only four vanes and four holes are shown, but more, e.g. six or eight, may be used. Such vanes and discs may be manufactured by pressing to form a sleeve which may be fitted and brazed or welded onto the jacket 5 with the intention that each gas mixing zone also serves to remove hear from the reaction gases leaving each reaction zone. Although the schematic drawing of the reactor shows each reaction zone and gas mixing zone of the same size, these may be designed to differ according to the amount of heat to be removed. Additionally, each reaction zone may incorporate different and/or different loadings, of catalyst. Each metal catalyst support monolith suitably has 300-400 cells per sq.in. and is coated with a washcoat comprising ~36wt% alumina solids of D50 of approx 5μm and D90 of approx 16μm, to deposit approx 3g/m3. After drying and firing, the coated monolith was impregnated with

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a platinum salt solution to result in 5% by wt of Pt deposited on the alumina. The resulting coated monolith was again dried and fired before being assembled into the reactor.

Initial studies indicate that a reformate containing 1-2% CO in admixture with H₂, CO₂, N₂, water vapour with a small proportion of unreacted reformer feedstock (e.g. CH₃OH, CH₄, gasoline), at 180°C may be converted to a feed gas suitable for a fuel cell, containing below 10pm CO, and at 140°C.

The reactor of the invention is compact and exhibits relatively low pressure drop. Manufacturing costs are relatively low. The reactor is believed to be suitable for mobile or automobile uses as well as stationery uses. It is presently envisaged that the reactor shell will be lagged to control heat loss and prevent injury. The reactor can be manufactured simply and inexpensively, using pressed and welded parts. It is desirably operated to achieve low pressure drop, and the particular embodiment illustrated is designed for a gas hourly space velocity of approx. 30,000h⁻¹ gas throughput.

The reactor as described may be altered in a variety of ways without departing from the central inventive concept.

The invention is now illustrated below in a working example of a two-stage reactor, fed with a synthetic reformate gas mixture containing N2, H2, CO, CO2 and H2O. Each of the two stages consists of a mixing zone with three off-set spiral vanes, followed by air injection accomplished through a number of apertures in an annular ring, producing a swirling, mixing gas. The thus mixed hydrogen and air mixture passes through an annular metal honeycomb support mounted on a central cooling jacket. The cooling jacket is fed counter-current to the flow of hydrogen with a heat transfer fluid, conveniently water under pressure, and the metal honeycomb support is mounted in good heat exchange contact with the cooling jacket. The metal honeycomb carries a selective oxidation catalyst. Gas may be extracted for analysis between the two stages and/or after the second stage. By setting the composition of the feedstock gas mixture according to that analysed after the first or second stage, the test two-stage reactor can model a three or four stage reactor.

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The detailed operating conditions and other experimental detail for one typical run of the above reactor now follow. It was remarkable that the pressure drop across the whole of the est reactor was about the measurement limit of the gauges being used, and was of the order of 1.5-3mbar (150-300Pa), at the two throughputs tested. This indicates that the reactor is very efficient, compact and requires very little energy to operate.

- Each catalyst support was an annular of internal diameter 3.85in (9.78cm), external
 diameter 5.85in (14,86cm) and 3.0in(7.62cm) length, made from "Fecralloy" steel
 and having 500 cells per sq in. Each support had been washcoated with alumina
 washcoat as used in exhaust gas catalytic convertors, followed by loading with 5%
 by wt of the alumina of Pt. Conventional manufacturing techniques were used.
- 2. The synthetic reformate had the compositions: 48%N₂(dry), 40%H₂ (dry), 10%CO₂(dry), 0.5-1.5%CO (dry) and 13% H₂O (of total flow). Two levels of CO concentration were chosen, a higher level to represent the output from autothermal reforming, and a lower level to represent the output from steam reformer or an optimised reformer, and two levels of throughput stated to be Nominal 3kWe (1kilowatt equivalent = 1,000 l/hr of H₂) and Nominal 6kWe, equal to 7,5000 l/hr of dry reformate and 15,000 l/hr of dry reformate respectively (8,620 l/hr and 17,240 l/hr of wet reformate).

It is plain from these results with a two-stage reformer that the compact and efficient reformer of the invention will be effective to remove CO from Hydrogen to any desired level, by conventional optimisation and especially by matching the number of stages to the requirements of the fuel cell system or other usage of the purified hydrogen.

TWO-STAGE ANNULAR SELOX REACTOR TEST RESULTS

	Sel**	969.0	0.640	0.556	0.684	0.582	0.579	0.540	0.505	0.472	0.562	0.433	0.520	0.503	0.428	0.502
GE	CO out	1610	740	950	1840	290	232	29	28	55	373	20	722	51	105	62
SECOND STAGE	Temp out, C	212	217	227	175	220	145	136	130	146	157	150	176	185	198	174
SEC	Temp in, C	167	167	169	129	191	138	127	121	131	134	138	148	140	146	176
	Air in, slm	2	3	4	3	3	0.5	8.0	0.1	1.2	1,0	0.1	-	2	3	0
	CO out %, dry	09.0	09.0	09:0	0.81	0.34	0.122	0.122	0.122	0.122	ć	٤	0.130	0.130	0.130	62ppm
	Temp out, C	212	214	210	149	215	176	167	191	172	166	179	167	162	166	202
贸	Temp in C	125	127	125	125	128	143	141	140	141	142	142	138	138	138	138
FIRST STAGE	Air in,	4	4	4	3	S	2	2	2	2	1.5	2.5	4	4	4	9
ír.	CO in % dry	1.5	1.5	1.5	2	1.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	Power kWe	3	3	3			,	6	3	-	-		9	9	9	9
	Ex No.	-	2	-	4		9	-	~		10	=	-	13	4	15

** Sel: (Selectivity) = CO consumed/2 x O_2 consumed

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CLAIMS

- A catalytic selective oxidation reactor, comprising a cylindrical reactor vessel (1) with counter current cooling means (5,6,7) and at least one stage, each stage being provided with an inlet for a first feedstock (2), and inlet for a second feedstock (8), gas mixing means (9) and a catalytic reaction zone (4), wherein the reaction zone comprises a selective oxidation catalyst deposited upon a metal support.
- A reactor according to claim1, wherein the metal support is mounted in good heat
 exchange contact with the cooling means.
 - A reactor according to any one of the preceding claims, wherein the metal support is a metal monolith.
 - A reactor according to any one of the preceding claims, wherein the cooling means comprises a central jacket mounted within the reactor.
 - A reactor according to claim 4, wherein the central jacket is connected via a coolant circuit to an external heat exchanger.
 - A reactor according to claim 4 or 5, wherein the central jacket is connected via a coolant circuit to a circulating pump.
- A reactor according to any one of the preceding claims, wherein the catalytic reaction
 zone(s) is/are annular and surround a central cooling means.
 - A reactor according to any one of the preceding claims, having from 2 to 10 stages.
 - A reactor according to claim 8, having four stages.
 - A reactor according to any one of the preceding claims, wherein the gas mixing means comprise annular mixing vanes or discs.

AMENDED SHEET

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- 11. A reactor according to any one of the preceding claims, connected to the output from a reformer such that CO is removed from the output gas to a level where the output from the reactor may be fed to a fuel cell.
- 5 12. A process for the removal of quantities of CO from a hydrogen feedstock by partial oxidation, comprising at least one step of admixing said feedstock with a quantity of oxygen, passing the resulting mixture over a selective oxidation zone comprising a selective oxidation catalyst on an annular metal support, and recovering a hydrogen product with reduced content of CO, whilst simultaneously cooling the selective oxidation zone by counter-current coolant flow.
 - A process according to claim 12, wherein the selective oxidation zone is maintained at a temperature in the range 100 to 250°C.
- 15 14. A process according to claim 12 or 13, providing hydrogen to a fuel cell.

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SUBSTITUTE SHEET (RULE 26)

Declaration and Power of Attorney For Patent Application English Language Declaration

As a below named inventor, I hereby declare that:

was filed on June 1, 2001 as

and was amended on June 1, 2001.

including the claims, as amended by any amendment referred to above.

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled

REACTOR AND PROCESS FOR REMOVAL OF CARBON MONOXIDE FROM HYDROGEN, the specification of which is attached hereto unless the following box is checked:

United States Application Number or PCT International Application Number 09/857,116

I hereby state that I have reviewed and understand the contents of the above identified specification.

I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR §

hereby claim foreign priority benefits under 35 U.S.C. §119(a)-(d) or § 365(b) of any foreign hereign patent or inventor's certificate, or § 365(a) of any PCT International application which

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		application for patent or inventor's	
		efore that of the application on which p	
Prior Foreign Appl	ication(s)	Pri	ority Not Claimed
9826222.3	Great Britain	<u>01 December 1998</u>	
(Number)	(Country)	(Day/Month/Year Filed)	
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(Number)	(Country)	(Day/Month/Year Filed)	
hereby claim the	e henefit under 35 U.S.C. (§ 119(e) of any United States provis	ional application(s)
listed below.	benefit under de d.c.o.	3 Tro(c) of unity offices elected provide	ionar appression(s)
(Application Number)	(Filing Date)		
(Application Number)	(Filing Date)		
I hereby claim the	benefit under 35 U.S.C. §	120 of any United States application(s	s), or 365(c) of any
PCT International	application designating the	United States, listed below and, inse	ofar as the subject
matter of each of	the claims of this applicat	tion is not disclosed in the prior Uni	ted States or PCT
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1.56 which became available between the filing date of the prior application and the national or PCT

international filing date of this application:

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	POWER OF ATTO agent(s) to prosecu connected therewith	ate this applic	named inventor, l cation and transact	hereby appoin all business in	nt the following atte the Patent and Tr	orney(s) and/or ademark Office				
	Allan Ratner Re Andrew L. Ney Re Kenneth N. Nigon Re Kevin R. Casey Re Benjamin E. Leace Re	eg. No. 23,031 eg. No. 19,717 eg. No. 20,300 eg. No. 31,549 eg. No. 32,117 eg. No. 33,412 eg. No. 24,842	Lawrence E. Ashery Christopher R. Lewis Robert L. Andersen Joshua L. Cohen Daniel N. Calder Louis W. Beardell, Jr. Jacques L. Etkowicz	Reg. No. 34,515 Reg. No. 36,201 Reg. No. 25,771 Reg. No. 38,040 Reg. No. 27,424 Reg. No. 40,506 Reg. No. 41,738	Jack J. Jankovitz Jonathan H. Spadt Christopher I. Halliday Scott A. Mckeown	Reg. No. 42,690 Reg. No. 45,122 Reg. No. 42,621 Reg. No. 42,866				
	Address all correspondence to: Christopher R. Lewis. Ratner & Prestia, Suite 301, One Westlakes, Berwyn, P.O. Box 980, Valley Forge, PA 19482-0980. Address all telephone calls to: Christopher R. Lewis at (610) 407-0700. Thereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements									
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l	by fine or imprisonn	nent or both	under Section 100	dements and	the like so made	are punishable				
	such willful false stat	tements may	jeopardize the validi	ity of the applica	ation or any patent is	ssued thereon.				
ŀ	Full name of sole or first in	nventor (given na	me, family name) Peter G	Seoffrev Grav						
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	Additional inventors	are being name	d on separately numbered	i sheets attached he	reto.					